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- Adhesive resin composition, laminate comprising this composition as adhesive layer, and process for preparation thereof.
- The styrene polymer resin composition comprising an ethylene/vinyl acetate copolymer, a styrene polymer resin, graft-modified polyethylene, a polystyrene elastomer and an ethylene/a-olefin copolymer in combination is disclosed. This adhesive resin composition shows an excellent adhesiveness between a metal and a thermoplastic resin. When this adhesive resin composition is used for bonding a polyester resin and a gas-barrier resin, a good bonding which is not degraded even by high-temperature heating can be attained.

# ADHESIVE RESIN COMPOSITION, LAMINATE COMPRISING THIS COMPOSITION AS ADHESIVE LAYER, AND PROCESS FOR PREPARATION THEREOF

## Background of the Invention

## 5 (1) Field of the Invention

The present invention relates to an adhesive resin composition and a laminate comprising this resin composition as an adhesive layer. More particularly, the present invention relates to an adhesive resin composition giving an excellent adhesiveness between a metal and a thermoplastic resin, and a metal/resin sheath laminate to be used for a laminate sheath cable, which is formed by using this resin composition. Furthermore, the present invention relates to an adhesive resin composition, of which the adhesive force is not reduced even under contact with high-temperature water, and a laminate having an excellent gas-barrier property and being capable of resisting a retort treatment, which is formed by using this adhesive resin composition.

(2) Description of the Related Art

An adhesive resin composition comprising three components, that is, a styrene resin, an ethylene/ninyl or acetate copolymer resin and a polyethylene resin graft-modified with an unsaturated carboxylic acid or a derivative thereof has been publicly known. This adhesive resin composition has an excellent adhesiveness between a metal and a thermoplastic resin and also between a polyester resin and an ethylene/vinyl acetate copolymer, which are used for a packaging material or the like to be subjected a retort treatment.

This adhesive resin composition is used for a laminate sheath cable having a resin sheath arranged on the outer side of a barrier material for a cable and a cable core, and since this laminate sheath cable is excellent in such characteristics as mechanical properties, corrosion resistance and molsture resistance, this cable is widely used at the present. In this laminate sheath cable, the above-mentioned adhesive resin composition is used for bonding a metal tape composed of aluminum, copper or the like, to be used as the barrier laver, to a sheath resin composed mainly of low-density opterfylene.

The laminate metal tape is generally prepared by extrusion lamination of a metal and a sheath resin or bonding a metal layer to a tusion-bonding resin layer film and a sheath resin layer film. In case of conventional fusion-bonding resins, streaks are often formed on the laminated fusion-bonding resins layer.

These streaks are formed because of local changes of the thickness of the fusion-bonding resin layer and the presence of these streaks degrades the adhesion between the metal and sheath resin and reduces the appearance characteristics of the product. Moreover, cracks are formed in a thin portion of the fusion-bonding resin layer and even the performances of the product are reduced.

In Japanese Unexamined Patent Publication No. 61:286044, we previously proposed a thermoplastic resin composition capable of preventing formation of streaks in the fusion-bonding resin layer, which comprises 97 to 45 parts by weight of an ethylene/vinyl acetate copolymer, 30 to 1 parts by weight of a styrene polymer resin, 15 to 1 parts by weight of polyethylene graft-modified with an unsaturated carboxylic acid or a derivative thereof and 30 to 1 parts by weight of a monowinyl aromatic hydrocarbon/folefin block copolymer. When a laminate metal tape is prepared by using this thermoplastic resin composition as the fusion-bonding resin, streaks are not formed in the fusion bonding resin layer, but the adhesiveness between the metal and a resin sheath layer is not completely satisfactory.

Accordingly, development of an adhesive resin composition which can prevent formation of streaks in the fusion-bonding (adhesive) resin layer and can give an excellent adhesiveness between a metal and a resin sheath layer is eagerly desired.

Although the above-mentioned adhesive resin composition is used for the production of a laminate of a polyester resin and a gas-barrier resin to be used for a packaging vessel or the lik and shows a good adhesiveness, if a heat treatment such a high-temperature filling treatment or a retort treatment is carried out at the production of this laminate or at the time of eating or drinking a content in the packaging vessel, the adhesive force is sometimes reduced by heating, resulting in peeling of the lay r and reduction of the gas permeation resistance.

As the adhesive resin composition capable of r taining a high adhesive force between layers of a

laminate even after a high-temperature treatment such as a high-temperature filling treatment or a retort treatment, we previously proposed in Japanese Unexamined Patent Publication No. 64-45445 an adhesive resin composition comprising (a) 95 to 50 % by weight of an ethylene/c-olefin copolymer having a melt flow rate of 0.1 to 50 g/10 min, a density of 0.850 to 0.900 g/cm³, an ethylene content of 75 to 95 mole% and an 35 X-ray crystallinity lower than 30%, (b) 5 to 50 % by weight of an ethylene/vinyl acetate copolymer having a melt flow rate of 0.1 to 50 g/10 min and a vinyl acetate content of 5 to 40% by weight, and (c) 1.0 to 30 % by weight, based on the sum of components (a) and (b), of partially or wholly graft-modified polyethylene having a grafting amount of an unsaturated carboxylic acid or a derivative thereof of 0.05 to 15 % by weight, a melt flow rate of 0.1 to 50 g/10 min, a density of 0.900 to 0.980 g/cm³ and an X-ray crystallinity of at least 30%, wherein the grafting ratio of the composition as a whole is 0.01 to 3 % by weight, the melt flow rate of the composition as a whole is 0.01 to 3 % by weight the melt flow rate of the composition as a whole is lower than 35 %. In this patent publication, we also proposed a laminate comprising a polyester or polycarbonate layer, an intermediate layer composed of this adhesive resin composition and a saponified olefin/vinyl acetate copylymer layer.

The above-mentioned achiesive resin composition has an excellent adhesiveness at normal temperature after a high-temperature treatment, but intertainiar peelling is sometimes caused in the above-mentioned laminate during a high-temperature filling treatment or a retort treatment.

Accordingly, development of an adhesive resin composition capable of completely preventing occurrence of interlaminar peeling in a laminate even under a high-temperature filling treatment or a retort to treatment is desired.

## Summary of the Invention

The present invention is to solve the above-mentioned problems of the conventional techniques, and it is a primary object of the present invention to provide an adhesive resin composition capable of preventing formation of streaks in a fusion-bonding resin layer and giving an excellent adhesiveness between a metal and a thermoplastic resin sheath layer, and a laminate formed by using this adhesive resin composition.

Another object of the present invention is to provide an adhesive resin composition having such an excellent heat-resistant adhesiveness at a high temperature that occurrence of interlaminar peeiing in a laminate can be completely prevented even during a severe treatment such as a high-temperature filling treatment or a retort treatment, while maintaining a practical adhesion strength at normal temperature after a high-temperature treatment.

Still another object of the present invention is to provide a laminate having an excellent gas permeation resistance and not causing interlaminar peeling at a high-temperature filling treatment or a retort treatment, which is formed by bonding a polycarbonate layer or polyalkylene terephthalate layer and a saponified ethylenevinyl acetate copolymer layer by using an adhesive resin composition as set forth above.

A first adhesive resin composition having an excellent adhesiveness between a metal and a thermoplas
tic resin sheath layer according to the present invention comprises (a) 86 to 45 parts by weight of a

ethylene/vinyl acetate copolymer, (b) 30 to 1 parts by weight of a styrene polymer resin, (c) 15 to 1 parts

by weight of polyethylene graft-modified with an unsaturated carboxylic acid or a derivative thereof, (d) to 1 parts by weight of a monovinyl aromatic hydrocarbon/olefin block copolymer elastomer and (e) an

ethylene/a-clefin copolymer, the total amount of components (a), (b), (c), (d) and (e) being 100 parts by

weight.

A first laminate formed by using the above-mentioned first adhesive resin composition according to the present invention comprises a layer of a metal such as aluminum, copper or iron, a layer of a thermoplastic resin such as a polyamide, a saponified ethylene/vinyl acetate copolymer, polyethylene or a polyester, and a layer of the first adhesive resin interposed between the two layers.

The second adhesive resin composition of the present invention, which has an excellent adhesiveness between a polyeater resin such as a polyalitylene terepithatiate or a polycarbonate and a gas-barrier resin such as a saponified ethylene/ninyl acetate copolymer, includes the following four embodiments.

An adhesive composition of the first embodiment comprises 100 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxlic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer being a styrene elastomer.

An adhesive resin composition of the second embodiment comprises 100 parts by weight of a soft power and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15 % by weight, the soft polymer

comprising (a) 20 to 100 % by weight of a styrene elastomer and (b) 80 to 0 % by weight of an ethylene/a-olefin copolymer having an ethylene content of 45 to 95 mole%.

An adhesive resin composition of the third embodiment comprises 100 parts by weight of a soft polymer and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer comprising (a) 20 to 100 % by weight of an ethylene/vinyl acetate copolymer having a vinyl acetate content of 5 to 40% by weight.

An adhesive resin composition of the fourth embodiment comprises 100 parts by weight of a soft polymer and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15 % by weight, the soft polymer comprising (a) 20 to 100 % by weight of a styrene elastomer, (b) 0 to 80 % by weight of an ethylene/α-olefin copolymer having an ethylene content of 45 to 95 mole% and (c) 0 to 80 % by weight of an ethylene/vinyl acetate copolymer having a vinyl acetate content of 5 to 40 % by weight.

The laminate of the present invention formed by the second adhesive resin composition includes the following two embodiments.

A laminate of the first embodiment comprises (I) a polyalkylene terephthalate resin layer, (II) an adhesive layer composed of a second adhesive resin composition as set forth above and (III) a saponified olefin/vinyl acetate copolymer layer.

A laminate of the second embodiment comprises (I) a polycarbonate resin layer, (II) an adhesive layer 20 composed of a second adhesive resin composition as set forth above and (III) a saponified olefin/vinyl acetate copolymer layer.

Laminates of the first and second embodiments are prepared by melting the respective resins independently in different extruders, feeding the melts to a die having a three-layer structure and co-extruding them so that the adhesive resin composition is interposed between the two other resin layers, or 25 by forming layers of two resins other than the adhesive resin composition in advance and melt-extruding the adhesive resin composition between the two layers.

# Detailed Description of the Invention

Adhesive resin compositions of the present invention, laminates formed by using these adhesive resin

compositions and processes for the preparation of these laminates will now be described.

(A) Adhesive Resin Compositions

# Ethylene/vinyl acetate copolymer

The ethylenevinyl acetate copolymer used in the present invention is a known ethylenevinyl acetate copolymer (EVA). In general, there is used an ethylene/vinyl acetate copolymer having a melt flow rate [MFR[E], ASTM D-1238, E] of 0.1 to 50 g/10 min, preferably 1 to 30 g/10 min, and a vinyl acetate conptent of 5 to 40 % by weight, preferably 8 to 11 % by weight in case of the first adhesive resin composition or 10 to 35 % by weight in case of the second adhesive resin composition. If an ethylene/vinyl acetate copolymer having MFR within the above-mentioned range is used, the melt viscosity is reduced, the moldability is improved and the adhesiveness is increased, and especially in the case of the second adhesive resin composition, the adhesiveness after a retort treatment can be further improved.

# Styrene resin

The styrene polymer resin used for the second adhesive resin composition of the present invention includes not only a homopolymer of styrene but also polymers and copolymers of styrene, nuclear substitution products thereof and substitution products of styrene formed by substitution at the α-position of the double bond, such as chlorostyrene, dichlorostyrene, methylstyrene, dimethylstyrene and α-methylstyrene. In general, in the present invention, a styrene polymer resin having a melt flow rate [MFR(G), ASTM D-1238, G) of 0.1 to 50 g/10 min, preferably 1 to 40 g/10 min, is used. If a styrene polymer resin

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having MFR (G) within the above-mentioned range is used, an adhesive resin composition having an excellent extrusion moldability is obtained.

# 5 Graft-modified polyethylene

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The graft-modified polyethylene used in the present invention is characterized in that the grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.01 to 15 % by weight, preferably 0.1 to 5 % by weight in case of the first adhesive resin composition or 0.1 to 10 % by weight in case of the second adhesive resin composition, the melt flow rate (ASTM D-1238, condition Elis 0.1 to 50 g/10 min, preferably 0.3 to 30 g/10 min, the density is 0.900 to 0.980 g/cm³, preferably 0.905 to 0.970 g/cm³, and the X-ray crystalinity is at least 30 %, preferably 35 to 75 %.

In this graft polyethylene, the polyethylene is partially or wholly graft-modified. The graft-modified polyethylene can be a product formed by graft-modifying an ethylene/a-olefin copolymer of ethylene with a minor amount, for example, up to 5 mole%, of at least one other a-olefin selected from propylene, 1-butene, 4-methyl-1-pentene, 1-bexene, 1-octene and 1-decene.

The graft-modified polyethylene is obtained by partially or wholly graft-modifying polyethylene or a thylene/-olefin copolymer with an unsaturated carboxylic acid or a derivative thereof. As the unsaturated carboxylic acid and its derivative, there can be mentioned, for example, unsaturated carboxylic acids such as acrylic acid, maleic acid, tumaric acid, tetrahydrophthalic acid, itaconic acid, citraconic acid, citraconic acid, corrotonic acid and Nacio acide (endocis-bicyclo[2,2]) hept-5-ene-2.3-dicarboxylic acid, derivatives thereof such as acid halides, amides, imides, anhydrides and esters. As specific examples of the derivative, there can be mentioned malenyl chloride, maleimide, maleic anhydride, citraconic anhydride, monomethyl maleate, dimethyl maleate, and glycidyl maleate, among these compounds, an unsaturated dicarboxylic acid or its anhydride is preferably used, and maleic acid. Nadic acid® and acid anhydrides thereof are especially preferably used.

For the production of a modification product by graft copolymerization of polyethylene with a grafting monomer selected from the above-mentioned unsaturated carboxylic acids and derivatives, various known processes can be adopted. For example, there can be adopted a process in which polyethylene is melted. 30 the grafting monomer is added to the melt and graft polymerization is carried out, and a process in which a solution of the grafting monomer in a solvent is added and graft polymerization is carried out. In each case, in order to increase the grafting efficiency of the grafting monomer, the reaction is preferably carried out in the presence of a radical initiator. The grafting reaction is generally carried out at a temperature of 60 to 350 °C. The amount used of the radical initiator is generally 0.001 to 1 part by weight per 100 parts by 35 weight of polyethylene. As the radical initiator, there can be mentioned organic peroxides and organic peresters such as benzoly peroxide, dichlorobenzoyl peroxide, dicumyl peroxide, di-tert-butyl peroxide, 2.5dimethyl-2,5-di(peroxidobenzoato)hexyne-3, 1,4-bis(tert-butylperoxy-isopropyl)benzene, laurovi peroxide, tert-butyl peracetate, 2,5-dimethyl-2,5-di(tert-butylperoxy)hexyne-3, 2,5-dimethyl-2,5-di(tert-butylperoxy)hexane, tert-butylperbenzoate, tert-butylperphenyl acetate, tert-butyl perisobutyrate, tert-butyl per-sec-oc-40 toate, tert-butyl perpiyalate and cumyl perpiyalate, and other azo compounds such as azobisisobutyrinitrile and dimethyl azoisobutyrate. Among these compounds, there are preferably used dialkyl peroxides such as di-tert-butylperoxide, 2.5-dimethyl- 2.5-di(tert-butylperoxy)-hexyne-3 and 1.4-bis(tert-butylperoxyisopropyl)benzene

## Monovinyl aromatic hydrocarbon/olefin black copolymer elastomer (styrene elastomer)

The monovinyl aromatic hydrocarbon/olefin block copolymer elastomer (styrene elastomer) is a polymer having a linear or branched block structure having a monovinyl aromatic hydrocarbon polymer block on at least one terminal thereof, which is represented by the general formula of

$$(A-B)_{n}, (A-B)_{n}-A' \text{ or } (A-B)_{n}-X$$

55 wherein A and A' represent a monovinyl aromatic hydrocarbon polymer block, B represents an olefin polymer block, n is an integer of from 1 to 5, m is an integer of from 2 to 7 and X represents a polyfunctional compound having a valency of m. Styrene and a-methylstyrene are preferably used as the monovinyl aromatic hydrocarbon, and styrene is especially preferably used. As the olefin, there can be

mentioned conjugated diolefins such as butadiene and isoprene, and a-olefins such as ethylene, propylene and 1-butene. The polymer block formed by polymerization of a conjugated diolefin can be hydrogenated. The block B may be composed of a copolymer of butadiene or isoprene with styrene or a-methylstyrene, so far as olefin units are contained in a major amount. In th. present invention, in the monovinyl aromatic s hydrocarbon/olefin block copolymer elastomer (d), the amount of the monovinyl aromatic hydrocarbon polymer blocks is generally 8 to 55% by weight and preferably 10 to 35% by weight a block copolymer having monovinyl aromatic hydrocarbon polymer blocks on both of the terminals is preferably used. These block copolymers are marketed, for example, under tradenames of Cariflex® TR and Kraton® G (each being a registered trade mark for a product supplied by Shell Chemicals).

## Ethylene/a-olefin copolymer

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The ethylene/c-olefin copolymer used in the present invention is an ethylene/c-olefin random copolymer which is characterized in that the melt flow rate [MFR(E), ASTM D-1238, condition E] is 0.1 to 50 g/10 min, preferably 0.3 to 30 g/10 min, the density is 0.850 to 0.800 g/cm³, preferably 0.850 to 0.890 g/cm³, the ethylene content is 75 to 95 mole%, preferably 75 to 90 mole%, in case of the second adhesive resin composition, or 45 to 95 mole%, preferably 45 to 90 mole%, in case of the second adhesive composition, and the X-ray crystallinity is lower than 30 %, preferably lower than 25%.

If an ethylene $\alpha$ -olefin copolymer having the above-mentioned characteristics is used, an adhesive resin composition having an excellent adhesiveness can be obtained. Especially, in the second adhesive resin composition, the adhesiveness after a retort treatment and the adhesiveness to a polyolefin are highly improved.

An ethylene α-olefin having 3 to 20 carbon atoms is used as the α-olefin constituting this ethylene/α25 olefin copolymer. As specific examples, there can be mentioned propylene, 1-butene, 1-hexene, 4-methyl-1pentnen, 1-octene, 1-decene, 1-letradecene and 1-octadecene. These α-olefin can be used alone or in the
form of a mixture of two or more of them.

The ethylene/a-olefin copolymer generally has a melting point (ASTM D-3418) lower than 100°C.

## Mixing ratios

In the first adhesive resin composition of the present invention, the ethylenevinyl acetate copolymer (a) is used in an amount of 96 to 45 parts by weight, preferably 85 to 50 parts by weight, the styrene polymer resin (b) is used in an amount of 30 to 1 parts by weight, preferably 25 to 5 parts by weight, the graft-modified polyethylene (c) is used in an amount of 15 to 1 parts by weight, preferably 10 to 2 parts by weight, the monovinyl aromatic hydrocarbon/olefin block copolymer elastomer (d) is used in amount of 20 to 1 parts by weight, preferably 18 to 3 parts by weight, and the ethylene/a-olefin copolymer (e) is used in an amount of 20 to 1 parts by weight, 18 to 3 parts by weight, per 100 parts by weight of the total amount of components (a) through (e).

In the second adhesive resin composition of the present invention, the graft-modified polyethylene is used in an amount of 1 to 30 parts by weight, preferably 2 to 28 parts by weight, per 100 parts by weight of the soft polymer.

In the first embodiment of the second adhesive resin composition, the soft polymer is a styrene 45 elastomer.

In the second embodiment of the second adhesive resin composition, a styrene elastomer and an ethyleneia-olefin copolymer are used in combination as the soft polymer. In this case, the soft polymer comprises 20 to 100 % by weight, preferably 20 to 90 % by weight, of the styrene elastomer, and 0 to 80 % by weight, of the styrene elastomer, and 0 to 80 % by weight, of the ethylene/a-olefin copolymer, with the proviso that the total amount of the styrene elastomer and the styrene/a-olefin copolymer is 100 % by weight.

In the third embodiment of the second adhesive resin composition, a styrene elastomer and an ethylene/vinyl acetate copolymer are used in combination as the soft polymer. In this case, the soft polymer comprises 20 to 100 % by weight, preferably 20 to 80 % by weight, of the styrene elastomer and 0 to 80 % by weight, preferably 10 to 80 % by weight, of the ethylene/vinyl acetate copolymer, with the proviso that the total amount of the styrene elastomer and the ethylene/vinyl acetate copolymer is 100 % by weight.

In the fourth embodiment of the second adhesive resin composition, an ethylene/c-olefin copolymer and an ethylene/vinyl acetate copolymer can be used together with the styrene elastomer as the soft polymer. In this case, the soft polymer comprises 20 to 100 % by weight, preferably 20 to 90 % by weight, of the

styrene elastomer, 0 to 80 % by weight, preferably 10 to 70 % by weight of the ethylene/a-olefin copolymer, and 0 to 80 % by weight, preferably 10 to 70 % by weight, of the ethylene/vinyl acetate copolymer, with the proviso that the total amount of the styrene elastomer, the ethylene/a-olefin copolymer and the ethylene/vinyl acetate copolymer is 100 % by weight.

The second adhesive resin composition of the present invention comprises the styrene elastomer and the graft-modified polyethylene as indispensable components, and the grafting ratio of the entire composition is 0.01 to 3 % by weight, preferably 0.05 to 2.5 % by weight. MFR of the total composition is 0.1 to 9/10 min, preferably 0.2 to 40 g/10 min, and the crystallinity of the total composition is lower than 35%.

The adhesive resin composition of the present invention is prepared by mixing the above-mentioned amounts of the above-mentioned components by known mixing means such as a Henschel mixer, a Y-type blender, a ribbon blender or a tumbling blender, or by melt-kneading the above mixture by a single-screw extruder, a twin-screw extruder, a kneader or a Banbury mixer and granulating or pulverizing the melt-kneader limiture.

Additives customarily used for thermoplastic resins, for example, a heat-resistant stabilizer, a weathering stabilizer, an antistatic agent, a lubricant, a slip agent, a nucleating agent, a dye or pigment and a plasticizer such as a hydrocarbon oil, can be added to the adhesive resin composition of the present invention, so far as the attainment of the objects of the present invention is not hindered.

# 20 (B) Laminates and Process for Preparation Thereof

In the laminate forming by using the first adhesive resin composition, the adhesive resin composition is interposed between a metal such as aluminum, copper or iron and a thermoplastic resin such as a polyamide, a saponified ethylene/vinyl acetate copolymer, polyethylene, a polycarbonate or a polyester.

This laminate can be prepared, for example, according to a process in which a film having a thickness of 10 to 200 µm is formed from the adhesive resin composition, the film is set between adherends, that is, the metal and thermoplastic resin, and fusion bonding is carried out to effect lamination, or a process in which the adhesive resin composition and the thermoplastic resin as the adhered are independently melted in different extruders and the melts are extruded through a multi-layer dye to effect lamination.

The laminate formed by using the second adhesive resin composition of the present invention comprises (I) a polyalkylene terephthalate resin or polycarbonate resin layer, (II) an adhesive layer composed of the second adhesive resin composition and (III) a saponified olefin/vinyl acetate copolymer layer.

The layer (I) constituting the laminate of the present invention is composed of a member selected from a polycarbonate resin,

The polyester resin used is a polyester comprising units of at least one dihydroxyl compound selected from aliphatic glycol, such as ethylene glycol, propylene glycol, 1,4-butanediol, neopentyl glycol and hexamethylene glycol, alicyclic glycols such as cyclohexanedimethanol and aromatic duhydroxyl compounds such as bisphenol, and units of at least one dicarboxylic acid compound selected from aromatic dicarboxylic acids such as terephthalic acid, isophthalic acid, adiple acid, sebacic acid and undecanedicarboxylic acid aliphatic dicarboxylic acids such as oxalic acid, succinic acid, adiple acid, sebacic acid and undecanedicarboxylic acid and allcyclic dicarboxylic acids such as hexahydroterphthalic acid. The polyester can be modified with a small amount of a polyhydroxyl compound or polycarboxylic acid having a valency of at least 3, such as a triol or a tricarboxylic acid, so far as the polyester shows thermoplasticproperties. As the thermoplastic oplyester, there can be mentioned polyethylene terephthalate, polybutylene terephthalate and a polyethylene is spothhalate/arendithalate conclumer.

The polycarbonate resin used in the present invention includes various polycarbonates and copolycarbonates obtained by reacting dihydroxyl compounds with phospene or diphenyl carbonates according to known processes. As specific examples of the dihydroxyl compound, there can be mentioned on hydroquinone, resorcinol, 4.4 -dihydroxydiphenylmetane, 4.4 -dihydroxydiphenylmetane, 4.4 -dihydroxydiphenylmetane, 4.4 -dihydroxydiphenylmetane, 4.4 -dihydroxydiphenyl-2.2-propane, 4.4 -dihydroxydiphenylmetane, 4.5 -dihydroxydiphenyl-2.2-propane, 4.4 -dihydroxydiphenylmetane, 4.5 -di

mechanical properties and transparency.

The above-mentioned adhesive resin composition is used for the adhesive layer (II) of the laminate of the present invention.

The layer (III) constituting the laminate of the present invention is composed of a saponified olefin/vinyl scattate copolymer. A saponified olefin/vinyl scattate copolymer prepared by saponifying an olefin/vinyl acetate copolymer having an olefin content of 15 to 60 mole%, preferably 25 to 50 mole%, to a saponification degree of at least 50 %, preferably at least 90 %. If the olefin content is within the above-mentioned range, thermal decomposition is hardly caused and melt forming can be easily performed, and the drawability, water resistance and gas permeation resistance are highly improved. If the saponification degree is higher than 50 %, the gas permeation resistance is highly improved.

As the olefin to be copolymerized with vinyl acetate, there can be mentioned ethylene, propylene, 1butene, 1-hexane, 4-methyl-1-pentene, 1-octane, 1-decene, 1 tetradecene and 1-octadecene. Among them, ethylene is especially preferably used in view of the mechanical stability and moldability.

For the preparation of the laminate of the present invention, there can be adopted, for example, a cosextrusion process in which the polyalkylene terephthalate resin or polycarbonate resin, the adhesive resin
composition and the seponified olefin/vinyl acetate copolymer are melted in different extruders, the melts
are supplied to a die having a three-layer structure and the melts are co-extruded so that the adhesive resin
composition forms an intermediate layer, or a sandwich lamination process in which the polyalkylene
terephthalate resin or polycarbonate resin and the saponified olefin/vinyl acetate copolymer are formed into
20 layers in advance and the adhesive resin composition is melt-extruded between the two layers. In view of
the interlaminer bonding force, the co-extrusion process is preferably adopted. The co-extrusion process
includes a T-die method using a flat die and an inflation method using a circular die. Either a single
manifold type using a black box or a multiple-manifold type can be used as the flat die. Known dies can
also be used in the inflation method.

The thickness of each layer of the laminates can be appropriately determined according to the intended use. In general, when the laminate is used as a sheet of film, it is preferred that the thickness of the polyalkyl terephthalate resin or polycarbonate resin layer be 0.02 to 5 mm, the thickness of the adhesive layer be 0.01 to 1 mm and the saponified olefin/vinyl acetate copolymer layer be 0.01 to 1 mm.

The laminate of the present invention can further have a structure of (I)/(II)/(II)/(II)/(I) in which layers (I) or aer arranged on both the sides, or a structure further comprising a polyolefin layer, for example, a structure of polypropylene/(II)/(III)/(II)/(II) or polyethylene/(II)/(III)/(II)/(II)/(II).

The present invention will now be described in detail with reference to the following examples that by no means limit the scope of the invention.

## Example 1

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High-density polyethylene (Hizex supplied by Mitsui Petrochemical Industries, Ltd.; MFR (E) = 5.5 g/10 min) was reacted with maleic anhydride to obtain graft-modified polyethylene having a maleic anhydride content of 0.5% by weight, MFR (E) of 3.0 g/10 min and a gel content lower than 0.1%.

To 5 parts by weight of the graft-modified polyethylene were added 60 parts by weight of an ethylene/vinyl acetate copolymer (vinyl acetate content = 10% by weight, MFR (E) = 9.0 g/10 min; hereinafter referred to as "EVA"), 20 parts by weight of polystyrene (supplied under tradename of "Denka Styrol GP200" by Denki Kagaku Kogyo, MFR (G) = 25 g/10 min; hereinafter referred to as "PS"), 10 parts by weight of a polybutdatiene block-hydrogenated polystyrene/polybutdatiene/polystylene block copolymer (supplied under tradename of "Krayton G1652" by Shell Chemicals, styrene content = 29% by weight) and 5 parts by weight of an ethylene/propylene copolymer [MFR (E) = 1.0 g/10 min, ethylene content = 80 mole%, X-ray crystallinity = 5%, density = 0.870 g/cm³), and the mixture was melt-kneaded and granulated by using an extruder provided with a Dulmage screw having a diameter of 40 mm to obtain a so composition (1).

The obtained composition (1) was melted at 200 and formed Into a press sheet having a thickness of 3 mm by using a compression molding machine. The physical properties (MFR and the density) of the press sheet were determined. The obtained results are shown in Table 1.

A fusion-bonding film having a thickness of 50 µm was formed from the composition (1) by using a smolding machine provided with a T-die having a diameter of 30 mm, and the presence or absence of streaks on the formed film was checked. By using this fusion-bonding film, an aluminum foll was bonded to a polyethylene sheet under conditions described below to obtain a laminate. (Structure)

A foil/film of composition (1)/polyethylene sheet

A foil: 200 µm in thickness, 10 in width and 15 cm in length

Composition (1) film: 50  $\mu$ m in thickness, 25 mm in width and 15 cm in length Polyethylene sheet: 2 mm in thickness, 25 mm in width and 15 cm in length

5 (Bonding Conditions) Temperature: 200 °C

Pressure: 6 kg/cm<sup>2</sup>

Time: 3 minutes

A test piece having a width of 10 mm and a length of 15 cm for measuring the bonding strength was cut out from the laminate by using a knife, and the 180 peel strength was measured at a pulling speed of

The obtained results are shown in Table 2.

## Example 2

To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 70 parts by weight of EVA, 10 parts by weight of FS, 10 parts by weight of Krayton G and 5 parts by weight of the ethylene/propylene copolymer, and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (2).

The physical properties (MFR and the density) of a press sheet of the obtained composition (2) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50 µm was formed from the composition (2) and the presence or absence of streaks was checked, and an aluminum foll/bookerthlene sheet laminate was obtained by using this film and the 1808 peel strength was measured.

The obtained results are shown in Table 2.

### Example 3

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To 10 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 60 parts by weight of EVA, 15 parts by weight of PS, 5 parts by weight of Krayton G and 10 parts by weight of an ethylene/propylene copolymer, and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (3).

The physical properties (MFR and the density) of a press sheet of the obtained composition (3) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50  $\mu$ m was formed from the composition (3) and the presence or absence of streaks was checked, and an aluminum foli/polyethylene sheet laminate was obtained by using this film and the 180 peel strength was measured.

The obtained results are shown in Table 2.

# Example 4

To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 65 parts by weight of EVA, 15 parts by weight of PS, 10 parts by weight of Krayton G and 5 parts by weight of an ethylene/butene copolymer [MFR (E) = 3.5 g/10 min, ethylene content = 85 mole%, crystallinity = 15%, density = 0.885 g/cm³], and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (4).

The physical properties (MFR and the density) of a press sheet of the obtained composition (4) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50 µm was formed from the composition (4) and the presence or absence of streaks was checked, and an aluminum foil/polyethylene sheet laminate was obtained by using this film and the 180° peel strength was measured.

The obtained results are shown in Table 2.

# Comparative Example 1

To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 80 parts by weight of EVA, 15 parts by weight of PS and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (5).

The physical properties (MFR and the density) of a press sheet of the obtained composition (5) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50 µm was formed from the composition (5) and the presence or absence of streaks was checked, and an aluminate foll/polyethylene sheet laminate was obtained by using this film and the 180° peel strength was measured.

The obtained results are shown in Table 2.

## Comparative Example 2

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To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were to added 70 parts by weight of EVA, 15 parts by weight of PS and 10 parts by weight of Krayton G, and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (6).

The physical properties (MFR and the density) of a press sheet of the obtained composition (6) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50  $\mu$ m was formed from the composition (6) and the presence or absence of streaks was checked, and an aluminum foil/polyethylene sheet laminate was obtained by using this film and the 180° peel strength was measured. The obtained results are shown in Table 2.

## Comparative Example 3

To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 70 parts by weight of EVA, 15 parts by weight of PS and 10 parts by weight of the ethylene/propylene copolymer, and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (7).

The physical properties (MFR and the density) of a press sheet of the obtained composition (7) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50 µm was formed from the composition (7) and the presence or absence of streaks was checked, and an aluminum follpolyethylene sheet laminate was obtained by using this film and the 180° peel strength was measured.

The obtained results are shown in Table 2.

## 40 Comparative Example 4

To 5 parts by weight of the maleic anhydride-grafted high-density polyethylene used in Example 1 were added 60 parts by weight of F2A, 15 parts by weight of F8, 10 parts by weight of high-density polyethylene [MFR (E) = 8.2 g/10 min, density = 0.965 g/cm³] crystallinity = 81%], and the mixture was melt-kneaded and granulated in the same manner as described in Example 1 to obtain a composition (8).

The physical properties (MFR and the density) of a press sheet of the obtained composition (8) are shown in Table 1.

In the same manner as described in Example 1, a fusion-bonding film having a thickness of 50 µm was formed from the composition (8) and the presence or absence of streaks was checked, and an aluminum so follipolyethylene sheet laminate was obtained by using this film and the 180° peel strength was measured.

The obtained results are shown in Table 2.

	Comparative	Example 4	6.5	-95		_	_			_			_		
		Exan	9	•	88		5			430			48	22	
	Comparative	Example 3	5.9	0.94	ı		96			510			43	49	
	Comparative	Example 2	5.2	0.94	ı		130			200			4	69	
	Comparative	Example 1	7.4	0.95	4		\$			190			48	72	
Table 1	Example 4		6.4	0.94	1		66			530			49	65	
Tat	Example 3		6.8	0.94	ı		100			490			46	63	
	Example 2		4.5	0.94	1		110			200			45	62	
	Example 1		5.2	0.94	1		26			490			48	9	
	Measurement	Method	ASTM D 1238	ASTM D 1505	ASTM D 638		ASTM D 638			ASTM D 638	,		<b>ASTM D 2240</b>	ASTM D 1525	
			MFR(E) (g/10min)	Density (g/cm³)	Strain at Yield	Point (kg/cm²)	Tensile Force at	Break Point	(kg/cm <sup>2</sup> )	Elongation	(between bench	marks) (%)	Shore Hardness(D)	Vicat Softening	Point ( C)

Table 2

0 0 0 0 × 0	5.2 4.9 5.2 5.1 5.1 3.0
0 0 ×	5.2 5.1 5.1
o ×	5.1 5.1
×	5.1
	***
0	3.0
×	4.0
0	2.5
	ļ
	0

# Example 5

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A 5-layer sheet was formed under conditions described below by using a composition (1) comprising 100 parts by weight of a styrene elastomer (Krayton G 1652 supplied by Shell Chemicals; hereinafter referred to as "SEBS") and 10 parts by weight of maleic anhydride-grafted modified polyethylene (MFR = 1.0 g/10 min, density = 0.925 g/cm<sup>3</sup>, crystallinity = 52%, butene content = 3.6 mole%, maleic anhydride 10 grafting amount = 1.0 g/100 g of polymer; hereinafter referred to as "MAH-PE-1"), a polycarbonate (Panlite L-1250 supplied by Teijin Kasei; hereinafter referred to as "PC"), a saponified ethylene/vinyl acetate copolymer (Kuraray Eval EP-F supplied by Kuraray, MFR = 1.3 g/10 min, density = 1.19 g/cm<sup>3</sup>, ethylene content = 32 mole%; hereinafter referred to as "EVOH"), and polypropylene (Hipol F 401 supplied by Mitsui Petrochemical Industries, Ltd.; hereinafter referred to as "PP"). 15 Sheet structure:

PC/(1)/EVOH/(1)/PP Layer thicknesses (µm):

80/50/50/50/80

Extruders:

20 40-mm screw diameter extruder, 260 °C (for PC)

30-mm screw diameter extruder, 250 °C (for (1))

30-mm screw diameter extruder, 210°C (for EVOH)

40-mm screw diameter extruder, 230°C (for PP)

With respect to the obtained 5-layer sheet, the interfacial bonding strength (FPC, g/15 mm) between the 25 layer of (1) and the EVOH layer and the interfacial bonding strength (FEVOH, g/15 mm) between the EVOH layer and the layer of (1) were measured at a peeling atmosphere temperature of 23 or 80°C and a peeling speed of 300 mm/min according to the T-peel test.

The sheet was subjected to a retort treatment at 131°C for 30 minutes. Then, the T-peel test was carried out under the same conditions as described above.

The obtained results are shown in Table 3.

### Example 6

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (2) comprising 75 parts by weight of SEBS, 25 parts by weight of an ethylene/propylene random copolymer (MFR = 1.0 g/10 min, ethylene content = 80 mole%, density = 0.865 g/cm<sup>3</sup>, crystallinity = 4%, hereinafter referred to as "EPR-1") and 10 parts by weight of MAH-PE-1 was used instead of the composition (1) used in Example 5.

The obtained results are shown in Table 3.

Then, a 5-layer sheet was formed by using the composition (2), the above-mentioned EVOH, polyethylene terephthalate (formed by adding a crystallization promoter to J135 supplied by Mitsui Pet; hereinafter referred to as "PET") and PP under conditions described below.

Sheet structure:

45 PET/(2)/EVOH/(2)/PP Laver thicknesses (µm):

80/50/50/50/80

Extruders:

40-mm screw diameter extruder, 280 °C (for PET)

50 30-mm screw diameter extruder, 250 °C (for (2))

30-mm screw diameter extruder, 210°C (for EVOH)

40-mm screw diameter extruder, 230 °C (for PP)

With respect to the obtained sheet, the interfacial bonding strength (FPET, g/15 mm) between the PET layer and the layer of (2) and the interfacial bonding strength (FEVOH, g/15 mm) between the EVOH layer 55 and the layer of (2) were measured under the same conditions as described above with respect to PC.

The obtained results are shown in Table 4

## Example 7

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 6 except that a composition (3) comprising 80 parts by weight of SEBS, 20 parts by weight of an s ethylene/vinyl acetate copolymer (MFR = 2.5 g/10 min, vinyl acetate content = 25% by weight; hereinafter referred to as "EVA-1") and 10 parts by weight of MAH-PE-1 was used instead of the composition (2) used in Example 6.

The obtained results are shown in Tables 3 and 4.

## Example 8

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A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 6 except that a composition (4) comprising 40 parts by weight of SEBS, 45 parts by weight of an 15 ethylener'i-butene random copolymer (MFR = 3.5 g/10 min, ethylene content = 89 mole%, density = 0.885 g/cm², crystallinity = 15%; hereinafter referred to as "EBR-1"), 15 parts by weight of EVA-1 and 20 parts by weight of MAH-PE-1 was used instead of the composition (2) used in Example 6.

The obtained results are shown in Tables 3 and 4.

#### Example 9

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (5) comprising 20 parts by weight of SEBS, 60 parts by weight of EBR-1, 20 parts by weight of EVA-1 and 10 parts by weight of MAH-PE-1 was used instead of the composition (1) used in Example 5.

The obtained results are shown in Table 3.

# 30 Example 10

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (6) comprising 60 parts by weight of SEBS, 20 parts by weight of EBR-1, 20 parts by weight of EVA-1 and 10 parts by weight of MAH-PE-1 was used instead of the composition (1) used in Example 5.

The obtained results are shown in Table 3.

# Comparative Example 5

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (7) comprising 100 parts by weight of EBR-1 and 10 parts by weight of MAH-PET was used instead of the composition (1) used in Example 5.

The obtained results are shown in Table 3.

## Comparative Example 6

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (8) comprising 85 parts by weight of EBR-1, 15 parts by weight of EVA-1 and 10 parts by weight of MAH-PE-1 was used instead of the composition (1) used in Example 5.

The obtained results are shown in Table 3.

#### 55 Comparative Example 7

A 5-layer sheet was obtained and the T-peel test was carried out in the same manner as described in Example 5 except that a composition (9) comprising 40 parts by weight of SEBS, 45 parts by weight of

EBR-1 and 15 parts by weight of EVA-1 was used instead of the composition (1) used in Example 5. The obtained results are shown in Table 3.

	-	dth)	23°C atmosphere after retort	350	1/90	1410	1110	200	410	1030	450	01
		(g/15 mmwidth)	80°C atmosphere	630	720	410	400	370	320	210	170	20
	ayer Sheet	FEVOH	23°C atmosphere	069	830	450	1150	520	390	1300	094	8
<b>~</b> 1	PC/EVOH Multi-Layer Sheet	ldth)	23°C atmosphere after retort treatment	390	860	750	290	810	570	110	1390	430
Table 3	PC,	(g/15 mmwidth)	80°C atmosphere	920	930	1010	044	350	029	180	120	610
		FPC	23°C atmosphere	046	1250	1190	860	069	, 830	910	1600	750
	rts by	a to a		100 10	75 25 10	80 10 10	పె <b>చ్</b> చి	2882	9889	90 00	£55	375
	Recipe (parts by	weight) of	Composition	SEBS MAH-PE-1	SEBS EPR-1 MAII-PE-1	SEBS EVA-1 MAII-PE-1	SEBS EBR-1 EVA-1 MAH-PE-1	SEBS EBR-1 GVA-1 MAH-PE-1	SEBS EBR-1 EVA-1 MAH-PE-1	EBR-1 MAII-PE-1	EBR-1 EVA-1 MAII-PE-1	SEBS EBR-1 EVA-1
				Example 5	Example 6	Example 7	Example 8	Example 9	Example 10	Comparative Example 5	Comparative Example 6	Comparative Example 7

Pecico (parts by weight) of Adhesive Resin Composition   PETITEVOH Multi-Layer Sheet   PETITEVOH (gri Smm width)   PETITEVOH (gr	ſ				l able 4	9.4			
FPET (g15mm width)   FPET (g15mm width)   FPUOH (g15mm sidth)		Recipe (parts by Adhesive Resin C	weight) of			PET/EVOH Mu	Iti-Layer Sheet		
SEBS         75         91 C atmosphere atmosph					FPET (g/15mm	width)		FEVOH (g/15mr	n width)
SEBS         75         960         980         980         980         970         700           MAHPE-I         10         320         730         460         420         420           SEBS         80         840         320         730         460         420           MAHPE-I         10         520         380         510         460         390           EBR-I         46         520         380         510         460         390           MAHPE-I         10         460         390         390         390			-	23°C atmosphere	. 80°C atmosphere	23°C atmosphere	23°C atmosphere	80°C atmosphere	23 C atmosphere after retort treatment
MAH-PE-1         10         320         730         460         420           SEBS         80         840         320         730         460         420           ENA-1         10         520         380         510         480         390           EBR-1         45         520         380         510         480         390           MAHPE-1         15         480         300         480         300         480	a)	SEBS EPR-1	মাম	096	880	088	810	002	800
SEBS ENA-1         80 20         840 840         320 320         730 320         460 460         420 460           SEBS EBR-1         45         520 460         380 510         510 460         460 380         380 510         460 460         380 380		MAH-PE-1	10						
EVA-1 20 MAH-PE-1 10 520 380 510 460 390 ENA-1 15 MAH-PE-1 20 MAH-PE-1 20 380 510 460 390 SI MAH-PE-1 20 SI MAH	-	SEBS	81	840	320	730	460	420	410
MAH-PE-1 10 520 380 510 460 380 ENA-1 15 MAH-PE-1 20 M		EVA-1	ଛ						
SEBS         40         520         380         510         460         390           EBR-1         45         52         380         510         460         390           MAHPE-1         15         7         7         7         7         7		MAH-PE-1	10						
EBR-1 EVA-1 MAH-PE-1		SEBS	위	250	380	510	460	390	430
EVA-1 MAH-PE-1	_	EBR-1	45						
	_	EVA-1	15						
	_	MAH-PE-1	8						

In Examples 5 through 10 and Comparative Examples 5 through 7, the crystallinity and density were measured according to the following procedures.

# (1) Preparation of Sample

The sheet was heated at 180°C by a hot press for 10 minutes and was rapidly cooled by a cooling press (water cooling) to prepare a sample.

## (2) Crystallinity

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The crystallinity of the sample obtained in (1) above was determined by the X-ray diffractometry.

## (3) Density

The density of the sample was measured at 23°C according to the density gradient tube method.

As is apparent from the results of the foregoing examples, since the first adhesive resin composition comprises (a) an ethylene/vinyl acetate copolymer, (b) a styrene polymer resin, (c) graft-modified polyethylene, (d) a monovinyl aromatic hydrocarbon/olefin block copolymer elastomer and (e) an ethylene/ $\alpha$ -olefin copolymer at a specific ratio, streaks are not formed at the extrusion molding, and a fusion-bonding (adhesive) film having a good appearance is obtained. Furthermore, the adhesive resin composition of the 25 present invention can give an excellent adhesiveness between a metal and a thermoplastic resin sheath

Accordingly, the adhesive resin composition of the present invention can be suitably used as a melt adhesive for laminates of laminate sheath cables and films of various packaging materials.

When the second adhesive composition of the present invention is used for bonding (I) a layer of a 30 polyalkylene terephthalate resin or polycarbonate resin and (III) a layer of a saponified olefin/vinyl acetate copolymer, the layers (I) and (III) can be so tightly bonded that peeling of the layers (I) and (III) is not caused even under high-temperature conditions as adopted for a high-temperature filling treatment or a retort treatment, and a practically sufficient bonding strength can be maintained at normal temperature after this high-temperature treatment.

Accordingly, a laminate comprising the layers (I) and (III) bonded by using the adhesive resin composition of the present invention has a high resistance to permeation of gases such as oxygen and therefore, the laminate has excellent properties as a retortable food packaging material.

#### 40 Claims

- 1. An adhesive resin composition, which comprises (a) 96 to 45 parts by weight of an ethylene/vinyl acetate copolymer, (b) 30 to 1 parts by weight of a styrene polymer resin, (c) 15 to 1 parts by weight of polyethylene graft-modified with an unsaturated carboxylic acid or a derivative thereof, (d) 20 to 1 parts by 45 weight of a monovinyl aromatic hydrocarbon/olefin block copolymer elastomer and (e) an ethylene/α-olefin copolymer, the total amount of components (a), (b), (c), (d) and (e) being 100 parts by weight.
  - 2. A laminate comprising an adhesive resin composition as set forth in claim 1, which is interposed between a metal and a thermoplastic resin.
- 3. An adhesive resin composition, which comprises 100 parts by weight of a soft polymer and 1 to 30 parts 50 by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer being a styrene elastomer. 4. An adhesive resin composition, which comprises 100 parts by weight of a soft polymer and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer comprising (a) 20 to 100% 55 by weight of a styrene elastomer and (b) 80 to 0% by weight of an ethylene/α-olefin copolymer having an ethylene content of 45 to 95 mole%.
  - 5. An adhesive resin composition, which comprises 100 parts by weight of a soft polymer and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated

carboxylic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer comprising (a) 20 to 100% by weight of a styrene elastorner and (c) 80 to 0% by weight of an ethylene/vinyl acetate copolymer having a vinyl acetate content of 5 to 40% by weight.

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- 6. An adhesive resin composition, which comprises 100 parts by weight of a soft polymer and 1 to 30 parts by weight of partially or wholly graft-modified polyethylene in which a grafting amount of an unsaturated carboxylic acid or a derivative thereof is 0.05 to 15% by weight, the soft polymer comprising (a) 20 to 100% by weight of a styrene elastomer, (b) 0 to 80% by weight of an ethylene/a-olefin copolymer having an ethylene content of 45 to 95 mole% and (c) 0 to 80% by weight of an ethylene/vinyl acetate copolymer having a vinyl acetate-content of 5 to 40% by weight
- 7. A laminate comprising (I) a polyalkylene terephthalate resin layer, (II) an adhesive layer composed of a resin composition as set forth in any of claims 3 through 6 and (III) a saponified olefin/vinyl acetate copolymer layer.
  - A laminate comprising (I) a polycarbonate resin layer, (II) an adhesive layer composed of a resin composition as set forth in any of claims 3 through 6 and (III) a saponified olefin/vinyl acetate copolymer layer.
  - 9. A method of producing a laminate of claim 2, which comprises forming a film or sheet of an adhesive resin composition, setting the resulting film or sheet between a metal and a thermoplastic film or sheet, and melt-adhering it.
  - 10. A method of producing a laminate of claim 7 or 8, which comprises melting (I) a polyalkylene terephthalate resin or a polycarbonate resin, (II) an adhesive resin composition and (III) a saponified product of an olefin/vinyl acetate copolymer, feeding them into a die of a multi-layer structure, and co-extruding them so that the adhesive resin composition is interposed between the polyester resin and the gas-barrier resin.
- 11. A method of producing a laminate of claim 7 or 8, which comprises forming (I) a layer of a polyalkylene terephthalate resin or a polycarbonate resin, and (III) an oblintivinyl acetate copolymer saponified product polyester resin layer, and then melt-extruding (II) an adhesive resin composition between the two layers.

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  Hoffmann, Eitle & Partner Patent- und
  Rechtsanwälte Arabellastrasse 4 Postfach
  81 04 20
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- Adhesive resin composition, laminate comprising this composition as adhesive layer, and process for preparation thereof.
- ② An adhesive resin composition comprising an ethylene/vinyl acetate copolymer, a styrene polymer resin, graft-modified polyethylene, a polystyrene elastomer and an ethylene/c-olefin copolymer in combination is disclosed. This adhesive resin composition shows an excellent adhesiveness between a metal and a thermoplastic resin. When this adhesive resin composition is used for bonding a polyester resin and a gas-barrier resin, a good bonding which is not degraded even by high-temperature heating can be attained.



# EUROPEAN SEARCH REPORT

Application Number

EP 90 11 5163

	DOCUMENTS CONSI	DERED TO BE RELEVAN	T	
Category	Citation of document with in of relevant pas		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL.5)
Y,D	WORLD PATENTS INDEX 87-040157, Derwent I London, GB; & JP-A-0 26-12-1986 * Abstract *	Publications Ltd,	1,2,5-	C 09 D 123/08 B 32 B 15/00 B 32 B 27/00 C 09 D 153/02
X			3-6	
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x	EP-A-0 188 901 (MI * Page 4, lines 2-2 8-30; page 6, lines 7-13; page 11, line	7; page 5, lines 5-13; page 8, lines	1-11	,
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x	al.)	MACHONIS, Jr. et	3-6	C 09 D B 32 B
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-	Place of search	Date of completion of the nearth	<del></del>	Example
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Y:p	CATEGORY OF CITED DOCUMI articularly relevant if taken alone articularly relevant if combined with a ocument of the same category echnological background on-written disclosure	E : earlier patent after the filing nother D : document cite L : document cite	document, but p date d in the applica I for other reas	rublished on, or tion ans



CL	AIMS INCURRING FEES
The overes	European patent application comprised at the time of filling more than ten claims.
	All claims fees have been paid within the prescribed time limit. The present European search report has been
	drawn up for all claims.
	Only part of the claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for the lirst sen claims end for those claims for which claims fees have been paid,
	namely claims:
П	No claims fees have been paid within the prescribed time limit. The present European search report has been
_	drawn up for the first len claims.
LA	CK OF UNITY OF INVENTION
	Division considers that the present European patent application does not comply with the requirament of unity of nd relates to several inventions or groups of inventions.
namely:	on naming on distance in Littletoning & Bi gabba an understraints
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	All further search less have been paid within the fixed time limit. The present European search report has
	boen drawn up for all claims.
	Only part of the further search fees have been pald within the lixed time limit. The present European search
	report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid.
}	namely claims;
	None of the further search fees has been paid within the fixed time limit. The present European search report
	has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims.
	namely claims:



# EUROPEAN SEARCH REPORT

Application Numb

EP 90 11 5163

	DOCUMENTS CONSIDE	RED TO BE RELEVANT	Γ	
Category	Citation of document with indica of relevant passage	ation, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
Х	of retowal passage WORLD PATENTS INDEX U- 86-064319, Derwent Put London, GB; & JP-A-61 22-01-1986 * Abstract *	ATEST, AN olications Ltd.	to claim	TECHNICAL FIELDS SEARCHED (tax CL3)
Τŧ	The present search report has been present search the present search the HAGUE CATEGORY OF CITED DOCUMENT.	Date of completion of the search 10-04-1992  S I: theory or princi	iple underlying	Examenses HMIDT H. R. the (aversion
Y:p:	CATEGORY OF CITED DOCCMENT: articularly relevant if taken alone articularly relevant if combined with anothocument of the same category choological background on-written disclosure	E : earlier parent d after the filing D : document cited L : document	ocument, but pr date i to the applicat for other reaso	ablished on, or



European Patent Office

# LACK OF UNITY OF INVENTION

The Search Division considers that the present European patent application does not comply with the requirement of unity of invention and relates to several inventions or groups of inventions.

#### namely:

- Claims 1,2,9 concerning EVA-based adhesive.
   Claims 5-8,10-11 partially as far as these deal with adhesive wherein the soft polymer contains mainly EVA.
- Claims 3,4 concerning styrene elastomer based adhesives Claims 5-8, 10-11 partially as far as these deal with styrene elastomer based adhesives.

